U.S. DEPARTMENT OF COMMERCE PATENT AND TRADEMARK OFFICE

## INFORMATION DISCLOSURE STATEMENT BY APPLICANT

(USE SEVERAL SHEETS IF NECESSARY)

ATTY. DOCKET NO. SUNESIS.002DV1

**APPLICANT** Wells et al.

FILING DATE

October 17, 2001

APPLICATION NO. 9/981547

**GROUP** 

1627

SHEET 1 OF 2

U.S. PATENT DOCUMENTS							
EXAMINER INITIAL		DOCUMENT NUMBER	DATE .	NAME	CLASS	SUBCLASS	FILING DATE (IF APPROPRIATE)
JE	1.	5,422,281	6/95	Harris et al.	436	501	
i	2.	5,571,681	11/5/96	Janda et al.	435	7.6	
	3.	5,783,384	7/21/98	Verdine	535	6	
	4.	5,958,702	9/28/99	Benner	435_	7.1	

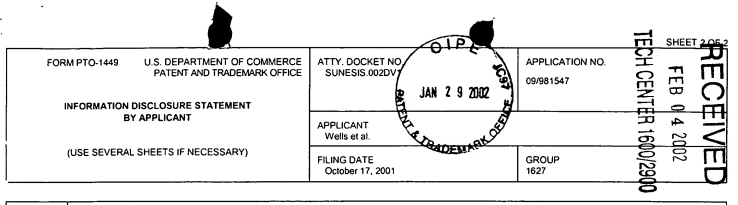
FOREIGN PATENT DOCUMENTS								
EXAMINER		DOCUMENT NUMBER	DATE	COUNTRY	CLASS	SUBCLASS	TRANSLATION	
INITIAL							YES	NO
THE	5.	EP 0 801 307	10/15/97	EP				
1	6.	WO 96/13613	5/9/96	PCT				
	7.	WO 96/27605	9/12/96	PCT				
	8.	WO 97/43302	11/20/97	PCT				
	9.	WO 98/11436	3/19/98	PCT		·	,	
d	10.	WO 98/11437	3/19/98	PCT				
								-

EXAMINER INITIAL	OTHER DOCUMENTS (INCLUDING AUTHOR, TITLE, DATE, PERTINENT PAGES, ETC.)
R	Abraham, D.J. et al., "How Allosteric Effectors Can Bind to the Same Protein Residue and Produce Opposite Shifts in the Allosteric Equilibrium" <u>Biochemistry</u> 34L150006-15020 (1995)

DATE CONSIDERED

\*EXAMINER: INITIAL IF CITATION CONSIDERED, WHETHER OR NOT CITATION IS IN CONFORMANCE WITH MPEP 609; DRAW LINE THROUGH CITATION IF NOT

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EXAMINER INITIAL	OTHER DOCUMENTS (INCLUDING AUTHOR, TITLE, DATE, PERTINENT PAGES, ETC.)					
JE	Boyiri, T. et al., "Bisaldehyde Allosteric Effectors as Molecular Ratchets and Probes" Biochemistry 34:15021-150 (1995)					
	13.	Bunyapaiboonsri et al., "Dynamic Deconvolution of a Pre-Equilibrated Dynamic Combinatorial Library of Acetylcholinesterase Inhibitors" ChemBioChem 2:438-444 (2001)				
	14.	DeJarias et al., "Use of X-ray Co-crystal Structures and Molecular Modeling to Design Potent and Selective Non-peptide Inhibitors of Cathepsin K" J. Am. Chem. Soc. 120(35):9114-9115 (1998)				
	15.	Erlanson et al., "Site-Directed ligand discovery" PNAS 97(17):9367-9372 (August 15, 2000)				
16. Foroozesh et al., "Aryl Acetylenes as Mechanism-Based Inhibitors of Cytochrome P450-Dependent Enzymes" Chem. Res. Toxicol. 10(1):91-102						
	17.	Hopkins et al., "Suicide Inhibitor of Cytochrome P450 1A1 and P450 2B1" <u>Biochem. Pharmacol.</u> 44(4):787-796 (1992)				
	18.	Lehn, Jean-Marie, "Dynamic Combinatorial Chemistry and Virtual Combinatorial Libraries" Chem. Eur. J. 5(9)2455-2463 (1999)				
	19.	Mathews et al., "N-Alkylaminobenzotriazoles as Isozyme-Selective Suicide Inhibitors of Rabbit Pulmonary Microsomal Cytochrome P-450" Mol. Pharmacol. 39(10):25-32 (1986)				
	-20,_	Misumi et al., "The p2 gag Peptide, AEAMSQVTNTATIM, Processed for HIC-1 Pr55 gag was found to be a Suicide Inhibitor of HIV-1 Protease" Biochem. Biophys. Res. Commun. 241(2):275-280				
	21.	Nicolaou et al., "Combinatorial Synthesis Through Disulfide Exchange: Discovery of Potent Psammaplin A Type Antibacterial Agents Active against Methicillin-Resistant Staphylociccus Aureus (MRSA)" Chem. Eur. J. 7(19):4280-4295 (2001)				
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	23.	Pollack, S. J. et al., "introduction of Nucelophines and Spectotroscopic Probes into Antibody Combining Sites" Science 242:1038-1040 (1988)				
	24.	Ramstrom and Lehn, "In Situ Generation and Screening of a Dynamic Combinatorial Carbohydrate Library against Concanavalin A" ChemBioChem 1:41-48 (2000)				
	25.	Woodcroft et al., "N-Aralkylated derivatives of 1-aminobenzotriazole as isozyme-selective mechanism-based inhibitors of guinea pig hepatic cytochrome P-450 dependent monooxygenase activity" Can J. Physiol. Pharmacol. 68(9):1278-1285 (1990)				
9	26.	Zhang et al., "Covalent Modification and Active Site-Directed Inactivation of a low Molecular Weight Phosphotyrosyl Protein Phosphatase" <u>Biochemistry</u> 31(6):1701-1711 (1992)				

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EXAMINER OF	DATE CONSIDERED 3/3/53
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